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(54) Self-tanning dihydroxyacetone formulations having improved stability and providing enhanced delivery

(57) A composition is provided which is useful for self-tanning skin coloring and is characterized by fin-incroved stability, which comprises from about 0.5% to about 20.0% by weight, based on total weight of said composition, of a self-taining skin coloring agent subject to chemical instability, which is preferably disydroxyacetone, from about 2.0% to about 4.00% by weight of a polyethoxyglycol, which is preferably emborydlycol, and from about 0.1% to about 15.0% by weight of a polyel comprising a polyhydric compound having at least three bydroxy groups and at least three

carbon atoms, which is preferably D-scribitol. The self-tanning composition may further optionally contain from about 0.1% to about 0.5% to yeeight of a water soluble dhydroxyl compound having at least two, and up to eight carbon atoms, which is preferably ethyriene glycoi; and the self-tanning composition may still further optionally contain an accidifying agent in amount sufficient to maintain the pH of said total composition at from about 3.5 to about 4.5, which is preferably sorbic acid. Ose-metologic products and methods of famining are also prometologic products and methods of famining are also prometologic products and methods of famining are also pro-

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Description

FIELD OF THE INVENTION

The present invention is in the broader technical field relating to compositions for the loquical application of active agents to the skin with sufficient perentation to be efficacious. Such active agents can be commeted agent as the relationship and the present invention is, further, in the narrower technical field relating to commet one commendation where the active agent is a unstress tenning compound, in particular disprogressions, which is delivered by tipical application of a commetic product formulation containing diright conjugations have sufficient stability and stail product in this narrower technical field; it is appreciated that desirable product formulations have sufficient stability and skin premittation to provide an even tan of the desired color in a timely product formulations have sufficient stability and skin premittation to provide an even tan of the desired color in a timely

BACKGROUND OF THE INVENTION

The prevailing cultural esthetic among fair-ekhned persons in many countries and places is to have a skin which is tanned, i.e., which has a light yellowish brown or even deeper brown color produced by exposure to the sun or to an artificial source of ultraviolet leight. The lanning of the skin is produced as a result of the darkening of preformed melanin, accelerated formation of new melanin, and retention of melanin in the epidemia as a result of relardation of kestarbization. The darkening of extesting melanin, possibly caused by oxidation and referred to as the Maircowsky phenomenon, begins within a few seconds of exposure to long-wave ultraviolet light and is complete within minutes up to a tew hours, depending on the individual involved.

For those many individuals who wish to achieve a tarned skin, the most readily available means for doing so is by exposure of their skin to natural suinight. However this method carries with it certain hazards, chief among which is the risk of surbmun, £e, actual liquip to the skin produced by excessive exposure to ultraviolet rays. The rijury is accompanied by enythema, tendemess, and sometimes bilatering. Furthermore, excessive exposure to ultraviolet are adiator is considered by the medical community to be a leading factor in the oncegoness of melanomas and other skin cancers, as well as an accelerating agent in the aging of the skin, particularly its tendency to sag and wrinkle, in order to mitigate or prevent such excessive exposure, such individuals will usually seek the protection of various sunscreen products, which contain sunscreen agents that act either by absorbing the ultraviolet radiation or by reflecting incident light. Sunscreen products also provide the user with other protections and benefits which are desirable, such as those evallable form skin moisturizine, prodective and healing compounds, anesthetic and anti-fulliammatory agents, etc. Certain embodiments of the present invention utilize such conventional sunscreen materials in order to schieve a product with the most universal appeal and with the greatest possible to make or most universal appeal and with the greatest possible to time universal sunscreen in an order to schieve a product with the most universal appeal and with the greatest possible to the universal to the universal and an incident of the protection of the

Where ultraviolet radiation is the source of skin tarning, it should be noted briefly that such electromagnetic radiation lies beyond the violet end of the spectrum, where its name. Ultraviolet radiation lies between violet rady and ended the violet rady is the special properties. Over 99% of ultraviolet radiation has wavelengths between 200 and 400 and is referred to as ultraviolet radiation has wavelengths between 320 and 400, and is referred to as ultraviolet and crimination. When the remaining rife of ultraviolet radiation comprises ultraviolet B or UVB radiation, which has wavelengths between 250 and 320 mm. UVB causes sunburn and a number of damaging photochemical changes within cells, including damage to DNI, kleading to premature aging of the skin, premalignant and malignant changes, and a variety of photocenstrivity reactions. Wavelengths between 200 and 290 mm characterize ultraviolet C radiation, or carriedy of photocenstrivity reactions. Wavelengths between 200 and 290 nm characterize ultraviolet C radiation, or carriedy of photocenstrivity reactions. Wavelengths between 200 and 290 nm characterize ultraviolet C radiation, or carriedy of photocenstrivity reactions. Wavelengths between 200 and 290 nm characterize ultraviolet C radiation, or carriedy of photocenstrivity reactions. Wavelengths between 200 and 290 nm characterize ultraviolet C radiation, or carriedy of wavelengths between 200 and 290 nm characterize ultraviolet C radiation, or carriedy to the carried until the earths surface. Many presents, in order to take advantage of naturally occurring sunlight, have resorted to the use of ultraviolet spith, which are also much easier to regulate with respect to the amount of ultraviolet radiation to which the skin is exposed, than is the case with natural sunlight. On the other hand, there are many persons who, to a variety of reasons, make use of so-called self-tenning or sunless product formulations in order to achieve

a satisfactory tam. For example, many persons have skin complexions which do not tan readily or evenly when exposed to sunlight, and many others suffer significant adverse side effects from sun exposure, including severe surbum. For such people, once a tan is obtained, there is a great deal of interest inmaintaining or extending the life of that tan, which will naturally dissipate over time as the layers of skin which actually comprise the tan become necrotic and are eventually slowled off. All such persons would greatly benefit from access to any means of obtaining a tanned skin, or at least a skin having the appearance of a naturally tanned skin, which did not require being exposed to sunlight, with all of the having the appearance of a naturally tanned skin, which did not require being exposed to sunlight, with all of the satient adverse effects which arise from such exposure to ultraviolet radiation. Accordingly, such persons in ever greater numbers have turned to so-called self-tanning or sunless product formulations in order to achieve a 'tan without greater numbers have turned to self-tanning products as a way of achieving a more affected by exposure to ultraviolet radiation, also have turned to self-tanning products as a way of achieving a more

uniformism, of obtaining a deeper tan with significantly less total exposure to sunlight, and of extending the natural lite of the tan which they have acquired for themselves. For such persons, a suitable product is to be found in those embodiments of the present invention which combine the components of a self-tanning composition with the components of a suscept formulation.

Salf-tanning agents and formulations of various types and compositions are known in the art. For example, sunless tanning agents which have been discovered and used herestopes include diphoroxyactore (DHA), globeraldehyde- (glycerol aldehyde) and related alcohol-aldehydes, various indoles and imidazoles and their derivatives, and pigmentation agents approved for use with humans, ag., methoscales and intoxactor. Dihydroxyacetore in particular is currently the self-anning agent most widely used, and it has been known in the art for some time as an articular is currently the self-anning agent most used be used herealter, the term "dhydroxyacetone" will be used to mean not only that specific corroposition of matter, but all of the other species described further herein which may be substituted for dhydroxyacetone in accordance with the objectives herein enumerated, whether with betteror poorer results as measured by the several criteria has further below described in more details.

It has been widely postulated in the art that dihydroxyacetone achieves this effect by reacting with various skin proteins and amino acids to produce a tan-cobred reaction product. Dhydroxyacetone has been used in a number of self-tanning formulations over the years combined, e.g., with colorants and sunscreen agents. Dihydroxyacetone, which may also be named as 1,3-dhydroxy2-propanone, has the following structural formula:

(1)

which corresponds to a elementary formula of $C_2 V_6 C_3$ and a molecular weight of \$0.08. Distrytoxyacetone is a crystalline powder which is fairly hygroscopic and has a melting point of about 75-80°C. It has a characteristic odor. The normal form in which dishydroxyacetone exists is a dimer, which is slowly soluble in 1 part water and 15 parts ethanol. However, when dishydroxyacetone is freshly prepared it reverts rapidly to the monomer form, which is very soluble in water, inhand, after part declaration.

As already mentioned, dihydroxyscetone is probably the most wisely used commercially of all self-tanning agents, and it is the key active ingredient in numerous self-tanning products which have achieved solid market success. The precise mechanism by which dihydroxyscetone causes the self-tanning reaction was studied by 80th at al. and reported in J. Soc. Cosmet Chem., 35, 265-272, August 1984. They studied the Maillard reaction of dihydroxyscetone with various amino acids tound naturally in the hydrosigid pelicies and first layers of the stratum comment of the skin. The Maillard reaction is one between the amino groups of amino acids, peptides or proteins with the "glycosidic" hydroxyl group of ougrar, ulmately resulting in the formation of brownippients. This reaction is also referred to commonly as the "browning" reaction, since this same reaction takes place when various foodstuffs are browned by heating during the process of their preparation for sellin.

However, despite such wide use and market acceptance, dihydroxyacotone suffers from several notable driawbacks which have both prevented products containing it from obtaining even wider market penetration, and precluded the production of a cosmetic product formulation with all of the optimum characteristics desired by the ultimate consumer. One of these notable drawbacks is an unacceptably short shell file obligatory drawprose prosesses as number of chemical and physical proporties which place significant inherent constraints on its use. Dhydroxyacotone in the monomer form is highly solution in water, which hereby exposes it to the action and affects of various agents, ransports in the many unintended environments, and makes its isolation difficult. The substantial oxidizing capacity of dhydroxyacotone, which is the utilizate mechanism by which it presumbly operates to produce a "lant", is a ensous problem tacing the cosmetic formulator or other antisan, since it can lead not only to undesired activity once applied to the skin of the utilimate consumers, but can also est of difficulties in terms of its interaction with the other components with which it is: a formulated Further, dihydroxyacotone does not possess a stable pH in solution, but is, rather, subject to a shifting pH with marker its formulation and utilizate use environment problematic to manage. Further still, driydroxyacotone is essocially sensitive to the degradative activities of bacters and other microorganisms, against the attacks of which it must be profected, both in its necessary.

A major rissult of the above-described properties of dhydroxyacetone is its degradation upon standing, which is likely to take place while the product containing it is in the possession of the ultimate consumer, if not even before purchase, while still in transit. This abbreviated shell life has required heretofere that products containing dhydroxy-actiona be used soon after preparation, and that said products be stored under conditions optimized with regard to temperature and relative humidity so that the progress of degradation could at least be delayed. Nevertheless, the proportion of dhydroxyacetone remaining active in each products but so been found to radiily diminish with time. In

addition to this loss of activity, the result of the chemical and physical degradation of dihydroxyacetone is the production of undesirable discoloration as well as the formation of disagreeable odors.

Another significant drawback of dryhdroxyacotone, which is not, however, related to its instability, is the inherent and unpleasant odor which results from the reaction of dhydroxyacotone and those components of the skin with which is comes in contact during the self-taming process. Accordingly, in order to address and overcome all of the above-described challenges posed by the properties of dhydroxyacotone, the arisan of ordinary skill in preparing cosmel and therapeutic products has been forced to exercise all of the formulation skills possessored by said artisan. The result, however, has not been the discovery of a totally successful product up until the time that the present invention was made as described further herein.

BRIEF DESCRIPTION OF THE RELEVANT ART

Protecting the skin from the adverse effects of exposure to ultraviolet radiation has long been of concern in the art partial with the present invention pertains. For example, Manox, U.S. Pst. 3, 019; 156; relates to a pharmaceutical proparation for oral administration which prevents, counteracts, or releves the effects on human skin and subcutaneous tissues of overapposure to sunlight. The preparation comprises paramichenezoes and (PARA) in combination with an arthitistamica operation and administration of the skin or exposure to sunlight. Further, Mejell U.S. Pst. 4,088,681 is concerned with compositions containing conjugated dienes in contribution with a topical addisc compatible surfected not containing carrier which are used to control the chronic effects.

of prolonged exposure to sunlight.

In addition to protecting the skin from exposure to ultraviolet radiation, the art has also been concerned with freating the skin for conditions which might be associated with such exposure, e.g., dryness, For example, Musher U.S. Pat. 4454.159 concerns proparations for treating irritated, prurities and dys skin which contain a combination of plaids/fipcids comprising glycerol tricites and orbiter glyceride oils of certain fatty acids, tocophard, squalene, collagen protein, a humactant, and acpropyl palmatic. Chausase U.S. Pat. 4 570.220 relates to a skin conditioning cosmetic composition having enhanced conditioning and protection against dryness, compatible with personal care, topstad drug and insect repellant compositions, comprising a paratheryl microlativar, i.e. dry-lentheno, and an empliciant which includes a polyhydric achoridor humactant, preferably glycenol, and a polyether derivative, preferably a copolymer of polyethylene glycol are protected and polypropytene glycol having a molecular weight of from 100 to 2000. Sun scene additives, among other active agents are mentioned as includable in the cometic composition. Also, Ser and Miguel U.S. Pat. 5,437,859 is concerned with contenting hydrating characteristics to coments or pharmaceutical solid filty products, e.g., it pistics, and discloses a process for the preparation of stable, anhydrous solid dispersions comprising preferably from 40 to 85% of a firstly body and preferably from 40 to 25% of a dispersed objylythric alcohol. The polyhytric achoch the polyhytric alcohol has from 20 to 8 carbon and tomor 20 to 6 flydroody functions and includes, e.g., ethylene glycol, glycerin and 1,2-propane.

diol. Other additives typical for such cosmetic formulations are also described. However, the present invention is concerned in particular with that segment of the art to which it pertains which has been concerned with providing artificial or self-tanning compositions as a means of protecting the skin. Such compositions, nevertheless, must overcome many of the constraints placed on the makeup of ordinary protective compositions, and have even been used in combination therewith, as is contemplated with the present invention. For example, Vanterberghe and Rosenbaum U.S. Pat. 3,940,477 relates to a class of aminated y dialdehyes which may be used in cosmetic compositions to produce artificial tanning of the skin. McShane U.S. Pat. 4,434,154 relates to an artificial tanning and ultraviolet screening cosmetic composition which is stable after prolonged storage comprising 2.5 to 7.5% by weight of total composition of dihydroxyacetone, octyl dimethyl p-aminobenzoic acid, water, oil and surfactant. The compositions are oil-in-water emulsions and the surfactant is a sodium (C8-C18) alkyl sulfate, e.g., sodium lauryl sulfate. Ziegler and Crotty U.S. Pat. 5,232,688 concerns a self-tanning cosmetic compositions containing an ahydroxy substituted ketone or aldehyde, preferably dihydroxyacetone, and a polyacrylamide preferably cross-linked and having a molecular weight of from 1000 to 5,000,000 for thickening, providing nonstreaking performance and improving low odor maintenance. Optionally, propylene glycol is advantageously incorporated in amounts of 15%, preferably between about 25 and 90%, in order to improve color intensity on the skin. Barnett and Lowry U.S. Pat. 5,268,166 relates to the application of a color cosmetic composition to the skin by means of electrostatic spraying in which artificial tanning materials containing dihydroxyacetone are included among the types of cosmetic compositions which may be utilized in this manner. Many of the additives used in these color cosmetic compositions function to make the compositions electrostatically sprayable, or to give the compositions certain desired properties when applied. Crotty and Ziegler U.S. Pat. 5,302,378 relates to a self-tanning composition containing, e.g., dihydroxyacetone in combination with an anionic silicone polyol such as dimethicone copolyol phosphate which prevents streaking, and at least 15% of propylene glycol to improve color intensity. Durand U.S. Pat. 5.458,872 is concerned with a method for protecting and stabilizing dihydroxyacetone which comprises enclosing at least a portion of the dihydroxyacetone in dimer form in a watertight composition which releases the dihydroxyacetone upon application. The watertight composition, which pro-

tects the diflydroxyacetore from water, existation and thermal effects, comprises a mixture of water inscluble polymers, ag, vinylidene polychloride, coopylomes of dimethylaminoethylmethactyptate, cellutose acetaler, nitrocaldusee, and ethyl cellutose, and optionally oilly fats in the form of a mixture of monopylocetides, diglycerides, and triglycerides of fatty acids having a chain length between C₉ and C₉. Lentini and Zacchino U.S. Pal. 5.51.43. The relates to self-tames consentic compositions compressing dighytroxyacetome and at least one operiodizatine, an anatural cyclic oligesecthande of lax (c.), seven (β) for eight (γ) glucose residues, cycloamylose and cycloglucans which are naturally occurring clathrates, by means of which the cosmicic composition is rendered more stable as evidenced by longer storage life, has a reduced odor under storage and use, and has a reduction in the odor caused by the reaction of the dihydroxyacetone with the skin. Tanner and Pobinson U.S. Pal. \$.5,14.437 relates to chydroxyacetone erificial tranning compositions having improved stability which contain from 0.025% to 5% of a sall selected from metablisulfite salts, suffice salts, and hydrogen suffite salts, and mixtures thereous as the salts integral agent.

The compositions of the present invention, in addition to the below-described essential components thereof, contain a number of additional ingredients which, while optional, are nevertheless advantageous to include for the beneficial properties which they confer on said compositions. Said incredients or similar additives have sometimes been employed heretofore in the art. For example, Walther and Stein, U.S. Pat. 3,281,374 concerns a process for the preparation of water in oil emulsions with the aid of polymeric higher fatty alcohols as the sole emulsifying agents, which results in a stable, creamy to pasty, colorless and odorless product able to absorb above 90% of water and rapidly absorbed by the skin. The monomers used to make the polymeric higher fatty alcohols have, e.g., from 12 to 24 carbon atoms and a terminal primary hydroxyl group. Lachampt et al. U.S. Pat. 3,489,690 is concerned with emulsifiers for preparing stable and irreversible water-in-oil emulsions for cosmetic use which provide hydration to the comeal layer of the skin as well as facilitating penetration of the emulsion products into the skin. The emulsifier is an oxypropylenated-oxyethylenated alcohol having a linear saturated alkyl radical of from 12 to 20 carbon atoms. A cosmetic formulation comprising a sunburn preventing cream is disclosed which contains an ultraviolet ray filter and an emulsifier of the formula: C18H27-[OC₃H₆]_{6.5}-[OCH₂CH₂]₂-OH. Dumoulin U.S. Pat. 3,975,294 is concerned with a surface active composition to be used as an emulsifier with diorganopolysiloxanes to form transparent micro-emulsions. Vanlerberghe and Sebag U.S. Pat. 4,303,639 relates to a class of 1,2-alkanediols which are colorless, odorless and miscible with oils conventionally used in cosmetic preparations, and which provide lubricity, ease of application, and good spreadability without an oily or greasy feel. Calvo et al. U.S. Pat. 4,466,955 concerns skin bleaching compositions based on hydroquinone using an anhydrous cosmetic carrier comprising polypropoxylated and polyethoxylated fatty ethers. Murray et al. U.S. Pat. 4,810,489 relates to high oil phase pharmaceutical vehicles in the form of an emulsion system comprising 35% to 65% oil phase, 1% to 10% of an alkylated polyvinylpyrrolidone copolymer, and an emulsifier.

The art has also explored combinations of self-tanning compositions and other active agents. For example, Kurz et al. U.S. Pat. S. 589, 460 relates to a skin-coloring propagation completing a pytropycathony compound, e.g., dithy-droxyacetone, and at least one colorant which adheres to the skin and is preferably an organic bye which adheres by physical, rather than chemical bonding, e.g., and solorant and exoin derivatives. Solvania are selected from short and long-thair monoatechols, polyalochols, allytene gytost, etc. Bloom and Deckner U.S. Pat. 5,514,178 reletes to pharmaceutical compositions for topical application hardy enhanced penateration through the skin comprising an active agent, including self-tanning agents, e.g., dihydroxyacetone, a high molecular weight crosslinked cationic polymer of callstylaminosityly acrystet and methacrystes and a vinylic monomer, a high or low hyprophic lipophilic blackner (HLB) non-ionic surfactant, e.g., stearic acid ethoxystated with 1 mole of ethylene oxide (steareth-1), and an alkoxystated ether, e.g., polyxorypropiene buyl ether, having about 14 propylene oxide untils incorporated in its structure.

Despite many previous attempts in the art to overcome the disadvantageous properties of dihydroxyacistone, as reflected in the disclosures of the above-discussed patents, none have been successful and none of them has arrived at the approach and the solution to these problems which comprise the present invention and which are described in more detail further below.

SUMMARY OF THE INVENTION

In its broadest aspects, there is provided in accordance with the present invention a composition comprising three components; (1) from about 0.5% is about 2.0 0% by weight, based on total weight of said composition, of a self-anning skin coloring agent subject to chemical instability; (2) from about 2.0% to about 4.0.0% by weight, based on total weight of said composition of a polyehroydypyck, preferably ethocytightyced, and (5) from about 0.4% to about 1.60 by weight, based on total weight of said composition, or a polyel comprising a polyhydric compound having at least three hydroxy proups and at least three action atoms. The polyel forming one of the three components to said composition comprises one or more members independently selected from the group consisting of 1.26-hexanetrio, isopropylidene glycerol, objects, and the components of the

There is further provided the above-described composition in which the self-stanning skin coloning agent subject to the previous previous previous previous provided by the previous p

In accordance with other narrower, but no less preferred aspects of the present invention there is provided a composition comprising four components: the above-described components (1) through (3), comprising a self-tanning skin coloring agent subject to chemical instability, most preferably dihydrospacetione; polyethoxyglycol, preferably enhoxydglycol; and a polyot, most preferably scribto, as above-described, and (4) a water soluble dihydroxyl (diol) compound having at least two, and up to eight carbon atoms, preferably from two to five carbon atoms, present in an amount of from about 0.1% to about 8.0%, most preferably from about 0.4% to about 2.0% by weight, based on total weight of said composition. Preferably, said water soluble dihydroxyl (diol) compound will be one or most members selected from the group consisting of ethylene glycot, definylene glycot, intertylene glycot, water soluble polyethylena glycot, propries glycot, propries glycot, and propries

Futher in accordance with still other narrower, but no less preferred aspects of the present invention there is provided a composition expensive provided a composition expensive provided a composition expensive provided and provided and provided provided

The present invention also provides the above-described composition as a cosmetologic product for application to the hair, nails or skin of a subject for the purpose of tenning, octoring and/or derkening the same, wherein the remaining portion of seld composition comprises a cosmetologically acceptable carrier. The cosmetologically acceptable carrier includes one or more member is independently selected from the group consisting of aciditying and latituding antibactional, antitungal and entiprotozcial agents, across longer lating antimicrobial agents including antibactional, antitungal and antiprotozcial agents; demanding preservatives, anticidants; buffering agents, chelating agents, coloring additives including dyes and pigments; dermaclogically active agents; buffering agents, chelating agents, coloring additives including entuils/riping and sillaring agents and entuils of additives including entuils/riping and sillaring entuils/riping additives including entuils/riping entuils/riping/r

In accordance with the present invention there is further provided a method of tanning, coloring or darkening the heir, nails and/or skin of a subject comprising applying thereto an amount of the above described cosmetologic product sufficient to tan, color, or derken said hair, nails and/or skin of said subject to which it is applied to the extent desired by said applicant.

In accordance with the present invention there is still further provided a method for preparing the above-described cosmetologic product wherein said product is an autous solution, comprising the steps, carried out sequentially or simultaneously, of 1) combining the water, dishydroxyacetore, ethoxydiglycot, polyci, optionally dishydroxy1 compound, and optionally acidlying agent components of said product, optionally logether with netalizing agents, sequestering apents, antimicrobial preservatives, and/or antioxidants which are desired, optionally with a solvent therefor; and there agents, and therefor a contraction enhancing agents, copionizing additives, sunsceren agents, and/or other demandoigically active interest, and therefor a contracting the production of the production of the contraction of t

There is also further provided a method for preparing the above-described cosmetologic product wherein said

product is an emulsion, inverse emulsion, or suspension, comprising the steps, carried out sequentially or simultaneously, described further below.

In accordance with the present invention there is provided a composition comprising:

(1) from about 0.5% to about 20.0% by weight, based on the total weight of said composition, of an α -hydroxy aldehyde or ketone of the formula:

(11)

wherein H^1 is H, CH_2OH , $CHOHCH_2OH$, CH(OH)CH(=O), $CH(OCH_3)CH(=O)$, $CH(NH_2)CH(=O)$, or CH(NH-Phenyl)CH(=O); and H^2 is H or CH_2OH ;

(2) from about 2.0% to about 40.0% by weight, based on the total weight of said composition of a polyethoxyglycol of the formula:

· (II

wherein n is an integer of from 2 to 6; and R3 is H, (C1-C6)alkyl, or phenyl;

- (3) from about 0.1% to about 15.0% by weight, based on the total weight of said composition, of a polyol comprising a polyhydric compound having at least three pricary groups and at least three carbon atoms, prieferably comprising one or more members independently selected from the above-numerated group of such polyols;
 - (4) from about 0.1% to about 8.0% by weight, based on the total weight of said composition, of a dihydroxyl compound having from two to eight carbon atoms, preferably comprising one or more members independently selected from the above-enumerated group of such dihydroxyl compounds, and
 - (5) an acidifying agent in amount sufficient to maintain the pH of the total composition at from about 3.5 to about 4.5

Still further in accordance with the present invention there is provided a preferred composition as above-described comprising:

- (1) from about 4.0% to about 6.0% by weight, based on the total weight of said composition, of dihydroxyacetone;
- (2) from about 14.0% to about 25.0% by weight, based on the total weight of said composition of ethoxydiglycol;
- (3) from about 0.5% to about 15% by weight, based on the total weight of said composition, of D-sorbitol;
- (4) from about 0.4% to about 2.0% by weight, based on the total weight of said composition, of propylene glycol
- and butylene glycot; and
 (5) from about 0.04% to about 0.20% by weight, based on the total weight of said composition, of sorbic acid.

The present invention also provides the above-described compositions as cosmatologic products for application to the hair, nails or skin of a subject for the purpose of tanning, coloring and/or darkening the same wherein the remaining portion of said composition comprises a cosmatologically acceptable carrier as above-described.

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides a composition comprising from about 0.5% to about 20.0% by weight, based on total weight of said composition, of a self-tanning skin obbring agent subject to chemical instability which is preferably an onlydroy allothydo or leations, and more prelenably is dihydroxyacetone. The composition of the present invention has improved stability with regard to the self-tanning skin octobring agent component, especially dihydroxyacetone in that the remaining key components of the composition serve to overcome the chemical instability to which the self-

tanning skin coloring agent is subject. These key components of the composition of the present invention are effective alone or in combination to diminish or even prevent degradation of the self-tanning skin coloring agent, especially divdrovayeacher component.

The self-tanning skin coloring component is always used in combination with a polyethoxyglycol, especially ethoxydlycol in the compositions of the present invention. The additional key component which is required at a minimum in said compositions in order to counteract the chemical instability of the self-tanning skin coloring component is a polycl. The polycl component of said composition is effective in this regard when used abone. However, the reversal of said chemical instability is even more pronounced when the polycl component is used together with the further key component comprising a diveryor compound. Moreover and supprisingly, said reversal of chemical instability is still further pronounced when the polycl and dihydroxyl compound components are used together with the still further key component comercian an additivity as sent.

The basic causes of the chemical instability of the self-tarning skin coloring agent, especially where said agent is dihydroxyacetone, are still unknown even after investigation of this phenomenan. It is known that when the self-tarning skin coloring component and the polysthoxyglycol component are mixed tegether with water, the pH of the formulation mixture is about 4.0, at least initially, however, inexplicably the pH of the formulation mixture scon begins to drop, i.e., the mixture becomes more acidic and the self-tanning skin coloring agent, especially dihydroxyacetore, becomes unstable. The formulation mixture is observed to disintegrate with the formation of organic acids and formalin, leading to the creation of disagneeable oclores and unacceptable colors in the formation mixture.

It is theorized that the key components of the compositions of the present invention result in Istalization of the hydroxy groups of the self-lanning skin cloring agent, especially diphydroxyacetom, making it more resistant to chemical attack. It is also known that dihydroxyacetoms occur naturally as a dimer, but that only the monomeric form has self-tenning skin coloring properties. It is also possible that the sublikation of the self-tenning skin coloring properties. It is also possible that the sublikation of the self-tenning skin coloring sent by the components. Regardless of the manner in which it is achieved however, this stallization mesult is achieved as whether the source of the degradation is based on temperature, hydrolysis, pH, microbal stack, or some combination of these flactors. Further, it has been clearened that in the stablized compositions of the present timent in that the self-tenning skin coloring agent gradually disappears over time, but that the formutation mixture retains its integrity and remains stabling.

The preferred self-tanning skin coloring agent, dihydroxyacetone, 1,3-dihydroxy-2-propanone, may be represented by the following general structural formula:

(1)

which corresponds to an elementary formula of C₀H₀O₅ Dihydroxyacetone is a crystalline powder which is fairly tygroscopic, and the normal form in which it exists is a dimer, which is slowly soluble in 1 part water and 15 parts ethanol. However, when dihydroxyacetone is freshly prepared it reverts rapidly to the monomer form, which is very soluble in water, ethanol, either and acetone. This high degree of water solubility is one of several notable drawbacks from which dihydroxyacetone suffers and which has precluded up until the present invention the preparation of accomment product formulation containing dihydroxyacetone which possesses all of the optimum characteristics desired by the ultimate consumer.

Thus, one of the serious disadvantages of ditydroxyacetone is an unacceptably short shell life. In fact, dihydroxyacetone is characterized by properties which place significant inherent constraints on its use. Dihydroxyacetone in the monomer form is highly soluble in water. While this is a desirable property with respect to preparing formulations with dihydroxyacetone, it also creates a problem because it thereby exposes dihydroxyacetone to the action and affects of various agents, transports it into many unintended environments, and makes its solicition difficult for extempt in the art to overcome this undesirable property of dihydroxyacetone is disclosed in Durand U.S. Pat. 5,458,972 in which a method for protecting and stabilizing dihydroxyacetone relies on encapsulating it in a waterlight composition comprising a water-inscubble polymer, e.g., ethyl cellulose.

Another serious problem facing the cosmetic formulator or other artisan attempting to devise a useful composition containing dihydroxyacetone is the substantial oxidizing capacity of dihydroxyacetone, since it can lead not only our undesired activity once applied to the skin of the ultimate consumer, but can also lead to difficulties in terms of its interaction with the other components with which its formulated. Dihydroxyacetone is also characterized by an unstable

pH in solution, and its shifting pH will way from as high as about 6.0 to as low as about 2.0. At the lower pH's especially, dihydroxyacetone becomes very unstable, which makes its formulation and ultimate use environment problemate or manage. The optimum pH for stability of dihydroxyacetone is about 4.0, accordingly the compositions of the present invention have been buffered to pH 4.7. Another source of instability in dihydroxyacetone is its particular sensitivity to the degradative activities of bacteria and other microorganisms. It is thus necessary to protect dihydroxyacetone against such attack, both at the time it is incorporated in a formulation for eventual use, as well as at the time when such use actually takes place.

As used herein, the form "dhydroxyacclone" is sometimes broadly inclusive, especially of structurally related compounds. However, the more appropriate generic expression for such compounds is "a hydroxy addetydes and kelones. Nevertheless, the present invertion also includes the use of structurally dissimilar compounds, which are included within the rubric "self-arianing skin coloring agents". All such agents are similarly useful in producing or including the artificial larning process in human skin, as described herein, and are thus contemplated to be within the scope of the present invention. Dhydroxyacchonic itself may be represented by the following general structural formular.

A number of other compounds are already known in the art as capable of producing or inducing the same artificial tanning process in human skin as is produced or induced by dihydroxyacetone. Some of these are structurally similar, to dihydroxyacetone and include, *inter* also, the following:

Accordingly, in a preferred embodiment of the present invention there is provided a composition comprising from about 0.5% to about 20.0% by weight, based on total weight of said composition, of an e-hydroxy aldehyde or ketone of the formula:

(11)

wherein \mathbb{R}^1 is \mathbb{H} , $\mathrm{CH}_2\mathrm{OH}$, $\mathrm{CHOHCH}_2\mathrm{OH}$, $\mathrm{CH(OH)CH(=O)}$, $\mathrm{CH(OCH}_3\mathrm{)CH(=O)}$, $\mathrm{CH(NH}_2\mathrm{)CH(=O)}$, or $\mathrm{CH(NH-Phe-nyl)CH(=O)}$; and \mathbb{R}^2 is \mathbb{H} or $\mathrm{CH}_2\mathrm{OH}$. The above formula represents dihydroxyacetone when \mathbb{R}^1 is \mathbb{H} and \mathbb{R}^2 is $\mathrm{CH}_2\mathrm{OH}$.

Preferably, the composition of the present invention will contain from about 1.0% to about 15.0% by weight, more preferably from about 2.0% to about 10.0% by weight, more preferably still from about 3.0% to about 8.0% by weight. and most preferably from about 4.0% to about 6.0% by weight, based on total weight of said composition, of a selftanning skin coloring agent, preferably an α-hydroxy aldehyde or ketone of the above formula, and most preferably dihydroxyacetone...

It is well known in the art that the closest homolog to dihydroxyacetone, methyl glyoxal, has been identified as a metabolite or degradation product of dihydroxyacetone which is responsible for the development of discoloration in formulations containing dihydroxyacetone which have not been stabilized. Methyl glyoxal is also responsible for the production of an undesirable orange pigmentation when dihydroxyacetone is applied to the skin and is not present in a stabilized form. Accordingly, methyl glyoxal is not one of the compounds structurally similar to dihydroxyacetone which is contemplated to be a component of the compositions of the present invention. Methyl glyoxal may be represented by the following formula:

Other self-tanning skin coloring agents which are structurally dissimilar to dihydroxyacetone but are still contemplated to be within the scope of the present invention with regard to the ability of the compositions of the present invention to improve their stability and enhance their efficiency, include, but are not limited to the following:

(XII)

It has been discovered in accordance with the present invention that, surprisingly, a significant and unexpected degree of stability can be conferred on cosmetological and pharmaceutical compositions containing self-tanning skin coloring agents, preferably a hydroxy aldehydes or ketones, most preferably dihydroxyacetone, when such compositions additionally contain from about 2.0% to about 40.0% by weight, based on total weight of said composition of a polyethoxyglycol, perferably ethoxydiglycol, i.e., 2-(2-ethoxyethoxy)ethanol or Carbito®. The term "ethoxydiglycol" is sometimes used herein as broadly inclusive of the polyethoxyglycols which in accordance with the present invention are components of a composition which confers improved stability and enhanced performance on the above-described self-tanning skin coloring agents, preferably α-hydroxy aldehydes and ketones. Said polyethoxyglycols are compounds which may be represented by the following general structural formula:

(111)

n is an integer of from 2 to 6, and R3 is H, (C1-C2) alkyl, or phenyl. In the case where n is 2 and R3 is H, the resulting compound is ethoxydiglycol, which may be represented by the following formula:

(III-a)

Preferably, the amount of polyethoxyglycol, preferably diethoxyglycol which will be present in the compositions of the present invention will be from about 6.0% to bout 5.0% by weight, more preferably from about 10.0% to about 9.0% by weight, more preferably still from about 12.0% to about 27% by weight, and most preferably from about 14.0% to about 25.0% by weight, based on total weight of asid composition.

The broadest embodiment of the present revention compress compositions which contain one key component in addition to the above described self-anning sike notioning agent and optivehoxygloot. Their searnial self-anning has coloring agent and optivehoxygloot. Their searnial self-anning sike notioning agent and optivehoxygloot. The searnial self-anning sike notioning agent and specification atoms. The upper limit of the number of carbon atoms which may be present is estermined by the water soubtility of such a candidate polyol and the compatibility of such a candidate polyol and the compatibility of such a candidate polyol and the compatibility of such a candidate polyol coming a part of said composition of the present invention is preferred that the polyol forming a part of said composition of the present invention is given of glored, polyonyellymen sorbificial, gyearin (glored), digyparin, enthintion, mannitol, xyliol, D and L-sorbifol, glucose, fructoes, galactose, mannose, sucree, bettoes, treatices, and insolit. The polyol component of the composition of the present invention is present in an anound of from about 0.1% to about 1.0% by weight, more preferably firm about 0.4% to about 1.0% by weight, more preferably firm about 0.4% to about 0.4% to about 3.0% by weight, more preferably still from about 0.4% to about 3.0% by weight, more preferably still from about 0.4% to about 3.0% by weight, more preferably still from about 0.4% to about 3.0% by weight, and most preferably from about 0.5% to about 3.0% by weight.

The next broadest embodiment of the present invention comprises compositions which contain one further key component in addition to the above described essential elements comprising a self-taming skin coloring agent, polyethoxyglycol, and polyot. That optional but desirable additional element of the compositions of the present invention as dishydroxyl compound. The term 'ditrydroxyl compound as used herein is intended to mean a dihydric or diol compound and the state of the composition of the present invention or diol compound as the less the carbon atoms and up to as many as light carbon atoms. It is more preterred that addidityrid or diol compounds not one or more members selected from the group consisting of eithylene glycot, diethylene glycot, water soluble polyethylene glycot, water soluble polyethylene glycot, water soluble polyethylene glycot, butylene glycot, (i.e. 1,2-butanediol, 1,3-butanediol, and 1,4-butanediol, pentanediols, and haxylene glycots including 1,6-bexanediol and 2-ethyl-1,3-bexanediol.

"The next broadest, i.e., the least broad embodiment of the present invention comprises compositions which contain a still further sty component in addition to the above-described essential elements comprising a self-tenning skin coloring agent, polyethoxyglycol, a polyol, and a dihydroxyl compound. That optional but desirable additional element of the compositions of the present invention is an additiny agent in amount sufficient to maintain the pit of the total composition at from about 3.5 preferably about 4.0. While use of an actifying agent in composition at the present invention, it is contemplated that such use is a preferred embodiment of the present invention. It is theroized that the use of an actifying agent incryoses the method of the present invention on ymorning the stabilization reaction with the self-tenning skin coloring agent, especially dihydroxyacetone forward, i.e., the equation of the reaction illustrated further above is moved further to the right, toward completing.

Acidying agents which may be suitable for use in expension, compositions of the present invention will be apparent to the aristan industreed by the present description. It is letterate, of course, to emptoy a mild organic acid such as aceilic, adiplic, anisic, bentzoic, boric, cathonic, cirrusmic, citric, diphosphomic, to emptoy a mild organic acid such as aceilic, adiplic, anisic, bentzoic, boric, cathonic, cirrusmic, citric, diphosphomic, to expensive, for a control organic province, active, active province, and control organic province, active, active province amount depends upon both the strength and the concentration of the acidifying agent utilized, but will generally be in an amount of from about 0.01% to about 0.50% to wight, more about 0.02% to about 0.20% by weight, acein about 0.02% to about 0.20% by weight, more about 0.02% to about 0.20% by weight, more about 0.02% to about 0.20% by weight, more about 0.04% to about 0.20% by weight, acein the total weight of said composition.

To recapitulate, the present invention in its broadest embodiment is concerned with a composition comprising from about 0.5% to about 20.0% by weight of a self-tanning skin coloring agent, preferably an a-hydroxy aldehyde or ketone, most preferably dihydroxyacetone; from about 2.0% to about 40.0% by weight or a polyethoxygleco, preferably dif-

ethoxyglycol, and from about 0.1% to about 1.5.0% by weight of a polyto comprising a polyhydric compound having at the hydroxyglycol groups and at least three above and ones, all of said weight percentages being based on the total weight of and composition. These novel compositions of the present invention may be used for any purpose, but are particularly well suited for threspecition and cosmetic applications. More particularly well these novel compositions of the present invention provide self-teanning skin coloring cosmetic product formulations with optimal properties of storage trabularly which afforces song shelf lie and freedom from unwanted discontration and disagreeable corors, enhanced penetration which provides a more rapid and uniform tenning process, as well as improved resulting color quality and remarks of the provides o

Accordingly, the present invention is especially concerned with providing a cosmetologic product for application to the hair, rails or skin of a subject for the purpose of tanning, coloring and/or darkning the same, comprising a composition and escribed immediately above wherein a remaining weight percent portion of said composition comprises a cosmetologically acceptable carrier. The individual components which logather make up such a cosmetologically acceptable carrier are numerous and varied, but are also well known to the artissan of carriang skill. Such an artissan is a person who has studied pharmaceutical. Medicinal, or cosmetological chemistry and has an advanced degree in one or more of those fields and at least five years of working research experience in discovering new and useful pharmaceutical or cosmetological formulations. The components pix referred to and their functions will be pointed out in turn in order that the compositions of the present invention and their uses may be made even more clear.

Vehicle: the vehicle, other referred to as the base for the coemetologically acceptable carrier, may be any fluid material example of delivering the other components of the composition to the skin with acceptable absorption components in the components into the skin. The commendation that skin with acceptable acrier may be in a number of different and components into the skin. The commendation put is in the form of a spray or mist, aerosolized by means of a propellant or by mechanical means using arribent air, the carrier may be a semi-solid cream, tolton, get or emitten vehicle intended to be applied to the surface of the skin and thereafter frozed into the deeper layers of the skin through the deplication of pressure and friction from rubbing, where the carrier and its components are absorbed by the skin, the carrier may be in the form of a logical such as as colution, emulsion or supersion in which the carrier is a solvent to the other components are the discontinuous phase; or the carrier may be but also includes inverse emulsions in which the carrier is the discontinuous phase; or the carrier may be in the form of a solid such as a powder applied by dusting or spraying, a solid sick to be rubbed against the skin to which a portion of the solid adhrence, or a mask applied as a flexible, formable material or as a fliquid or gle which hardens to a solid by a process of solvent evaporation or crosslinking induced by heat, ultraviolic light for some other catalaying agency:

The preferred cosmelologically acceptable carriers for use in the present invention are an aqueous solution in which a suitable viscosity has been achieved, or an oil-in-water emulsion in which the continuous aqueous phase contains those active agents which are water soluble, and the discontinuous oil phase contains those active agents of the overall composition which are oil soluble. The emulsion inverts upon application to the lipophilic surface of the skin, whereby the active agents are absorbed into the skin form both the continuous and discontinuous phases. Where the carrier is an aqueous solution, the water acting as solvent is preferably delonized water which contains inorganic lone in concentrations insufficient to interfere with the stability or functioning of the overall composition, ag., the dihydroxyacetone, althoxydisycd, and sorbid are sufficiently water soluble to provide the acqueous solution, or sets are insufficiently water soluble to be added intently to the augueous solution, or sets are insufficiently water soluble and are first dissolved in water missoible solvents in which they are sufficiently solution, ag., propylene and buytene glycol, after which the water missible solvents in which they are sufficiently solution.

miscuise savent southern a south of the superconstance. Where the commeloopically acceptable carrier is an oith-n-water emulsion, an oil or lipid phase is established by the addition of hydrocarbon and/or silicone solvents together with various emulsifying agents and ast to permit stable constitations of the continuous and discontinuous phases. Whether aqueous solution or emulsion, the amount of water or aqueous carrier to be included in the compositions of the present invention will vary depending upon the desired consistency of the final product. By varying the amount of water, gelling agent, and/or surfactant present, it is possible to formulate a thick-flowing liquid or lotion, a semi-liquid thick cream, a pasta, a slick, a gel, an abcholic hydrogel, an emulsion, an alcoholic solution, or a formulation suitable for use in an aerosol. A consistent requirement of any such formulation, however, is that it can be, uniformly spread on the skin and dissorbed.

Into the vehicle of the cosmetologically acceptable carrier is introduced the desired pharmaceutically acceptable and cosmetologically acceptable additives whose compositions and functions are described in the below paragraphs.
Acidifying and Alkalizing Adams's are added to the composition largely to obtain a desired or predetermined plt.
Acidifying agents include, e.g., acetic acid, glacial acetic acid, maltic acid, and proportic acid. Stringer acids such as
hydrochoric acid, rintic acid and suffure acid should be avoided. Alkalizing agents include, e.g., edeol, potassium

carbonate, potassium hydroxide, sodium borate, sodium carbonate, and sodium hydroxide. However, alkalizing agents which contain active amine groups, such as diethanolamine and trolamine, not be used.

Acrosol Propellants - are required where the composition is to be delivered as a aerosol under significant pressure, and include, e.g., acceptable heliopenated hydrocators such as disholonofillusomenthane, cilchrotreritationosthane, and trichbromonofilusoromethane; nitrogen, or a volatile hydrocarbon such as butane, propane, isobutane or mixtures these?

Antimicrobial Agents Including Antibiacierial, Antifungal and Antiorotozcal Agents - are added where the compaint on will be applied to areas of the skin which are likely to have suffered adverse conditions or sustained abrazions or cuts which expose the skin to infection by bacteria, fungi or protozoa, and include, e.g., antimicrobial agents such as benzyl alcohol, chlorobutanol, phenylethyl alcohol, phenyletercuric acetale, potassium sorbate, and sorbic acid, and antifungal agents such as benzoic acid, butylerarben, ethylparaben, propyleparaben, and sodium.

In a preferred embodiment of the present invention, bency alcohol in combination with methylegaraben is used as the antimicrobial salegart. The combination of these agents provides an antimicrobial action which is gentile and does not interfere with the actions of the other components of the compositions of the present invention. The combination of bencyl actional dam dehtyleparaben plays a normal role, if any, in the stabilization of the self-anning skin coloring agents, especially dihydroxyacetone, used in the compositions of the present invention. By contrast, antimicrobial agents often found in self-anning commercial formulations comprise water-soluble compositions with very large atoms. However, it has been discovered that such compositions of the trobletome because they undergo a Maillard reaction with the dihydroxyacetons to form different acts products with have undesiráble properties. Although sorbic acid may also be employed as an ainthinicrobial agent, its preferred use, as discussed further above, is as an acidifying agent in accordance with the present invention.

Antimicrobial Preservatives - are added to the compositions of the présent invention in order to protect them against the growth of potentially harmful microorganisms, which susually invende the aqueous phase, but in some cases can also grow in the oil phase. Thus, preservatives with both aqueous and ligid solubility are desirable. Suitable antimicrobial preservatives include, e.g. aliqui besters of phydroxybarizok acid, propipates satis, phenoxybarizon, lamply preservatives include, e.g. aliqui besters of phydroxybarizok acid, propipates satis, phenoxybarizon, lamply paraben sodium, sodium dehydroxestate, benzialkonium chloride, benzelthonium chloride, and benzyl acholo. It is essential, of course, to select the preservative with a view toward possible incompabilities between the preservative and other ingredients of the composition. For example, antimicrobial preservatives which contain active armine groups are in a suitable: These include, e.g., hydration didrivities, quatemany ammonium compounds and cationic polymers, imidazolidinyl urea: diazolidinyl urea; and trisodium ethylenocliamine tetracetate (EDTA). Preservatives are preferably emolyced in amounts ranabine from about 0.01% to about 2.0% by weight of the foliat composition.

Antoxidents: protect all of the ingredients of the composition from damage or degradation by oxidizing agents present in the composition is self, the use environment, or the skin to which the composition is being applied, the use environment, or the skin to which the composition is being applied, in size of include, or e.g., anoxomer, ascorbly pairhitate, butylated hydroxyanisola, butylated hydroxylatena, hypophosphorous acid, potession materialswife, propie overly and dozdery allatels, sodium metabusities, suffur metabusities, suffur oxidios, and toccophosities.

<u>Buffering Agents</u> - are typically used to maintain a desired pH of a composition once established, from the effects outside agents and shifting equilities of componishes of the composition. However, one of the advantages of the compositions of the present invention is that they are self-buffering and seldom require the addition of a buffering agent in order to maintain the desired PK evertheless, in those rare instances where a buffering agent is added, if may be selected from among those familiar to the artisan skilled in the preparation of cosmelic formulation, e.g., calcium actitate, potassium metaphosphate, potassium prosphate monobasis, and trains' add.

<u>Chelating Agents</u> - help maintain the ionic strength of the composition and also serve a protective function in binding to and effectively removing a number of destructive compounds and metals, and include, e.g., edetate dipotassium, edetate discotium, and adulte acid.

Coloring Additives Including Dyes and Pigments. are especially useful adjuncts in the compositions of the present invention for the purpose of enhancing, Ioring, noreceing the intensity, and changing the faming color produced by the self-tanning agent, and include, e.g., caramet, 9-phenyl derivatives of 9-H-xanthen which are fluorescein derivatives of the formula:

1

(XIII)

where X^1 to X^4 are each independently H, NO_2 or halo; M is H or an alkali metal; and Ar is phenyl substituted by a metal carboxylate and optionally up to four halogen atoms; furobenzopyranone derivatives of the formulas:

Pigments must be chosen carefully to avoid unwanted catalysis of undesired side reactions in the compositions of the present invention. Thus, it would normally be unacceptable to select a metal coids or other active sets such as black fron coids, chromium coidse, yellow and red iron coidse, titanium dioxide, ultramarines (aluminosilicate black) and certain organic dyes such as cochineal carmine, azo dyes, and antiraquinone dyes; natural, colored cosmocially active substances which mask the hus produced by diffyactoxyacetions or other self-taming agent, such as the blue azulenes gualazulene and chamruzulene, the yellow flavanoids rulin and quercetin, enythrosin, bengal rose, orbitoxin, cyanonic pathninin, escini G, cosen 108, and Add Red 51.

Dermatologically Active Agents - are any one or more of the numerous and diverse known therapeutic agents applied to the skin for a dermatologically beneficial purpose, which include, e.g., wound healing agents such as peptide derivatives, yeast, panthenol, hexylresorcinol, phenol, tetracycline hydrochlonde, lemin and kinetin; glucocorticosteroids for treating inflammation, e.g., hydrocortisone, dexamethasone, betamethasone, triamcinolone, fluorinolone and methylprednisolone, retinoids for treating acne, psoriasis, cutaneous aging, and skin cancer, e.g., retinol, tretinoin, isotretinoin, etretinate, acitratin, and arotinoid, immunosuppressive agents for treating inflammation, e.g., dapsone and sulfasalazine; antibacterial agents for treating mild acne, e.g., resorcinol, salicylic acid, benzoyl peroxide, erythromycinbenzoyl peroxide, erythromycin, and clindamycin, and for treating impetigo, e.g., mupirocin; antifungal agents for treating tinea corporis, tinea pedis, candidiasis and tinea versicolor, e.g., griseofulvin, azoles such as miconazole, econazole, itraconazole, fluconazole, and ketoconazole, and allylamines such as naftifine and terfinafine; antiviral agents for treating cutaneous herpes simplex, herpes zoster, and chickenpox, e.g., acyclovir, famciclovir, and valacyclovir, antihistamines for treating prunitis, atopic and contact dermatitis, and psoriasis, e.g., diphenhydramine, terfenadine, asternizole, loratadine, cetirizine, acrivastine, and temelastine; topical anesthetics for rerlieving pain, irritation and itching, e. g., benzocaine, lidocaine, dibucaine, and pramoxine hydrochloride; topical analgesics for relieving pain and inflammation, e.g., methyl salicylate, camphor, menthol, and resorcinol; topical antiseptics for preventing injection, e.g., benza-Ikonium chloride and povidone-lodine; vitamins and derivatives thereof such as tocopherol, tocopherol acetate, retinoic acid and retinol.

Dispersing and Suspending Agents - include, e.g., poligeerian, povidone, and silicon dioxide, although povidone may cause Maillard side reactions and should be used with caution.

<u>Emollishs</u> - are agents, preferably non-oily and water-soluble, which soften and sooth the skin, especially skin, that has become dry because of excessive loss of water, and include, *e.g.*, yield carbon oils and waters such as mineral oil, petrolatum, microcrystaline was, and polyethylene; trighyeride series such as those of caster oil, cooks butter safflower oil, cotton each oil, corn cold, corn oil, olive oil, cod liver oil, atmond oil, avocado oil, paim oil, assame oil, squalene, and soybean oil, acelylated monoplycerides; entoxylated dypendes such as ethoxylated giveryer immonsterate, methyl, isopropyl, butyl and other allyl esters of C₁₀C₂₀ fatty acids, such as heayl learure, leorexylpalminate, decyl oiset, hexadecyl stearrie, isopropyl soctearante, dilscheryly adaptate, discoproryl sebacter, launyl lactate, myristyl lactate and cycl lactate, alkenyl esters of C₁₀C₂₀ fatty acids, such as oleyl myristate, clory isoprarier, locit, indied, crip-oile, fatty acids, such as pelargonic, launc, myrister, paramic, stearic, searier, isoprarier, oiler, lindied, crip-oile, arachidic, behane, and druce acids, clory C₁₀ of acids acids, such as launyl, myristyl, cetyl, hexadecyl, searyl, sostearyl,

hydroxystearyl, cleyl, lincleic, ricincleyl, behenyl, erucyl, and 2-octyl dodecanol alcohols; fatty alcohol ethers which are ethoxylated C10-C20 fatty alcohols such as the lauryl, cetyl, stearyl, isostearyl, oleyl, and cholesterol alcohols having attached thereto from 1 to 50 ethylene oxide groups or 1 to 50 propylene oxide groups; ether-esters such as fatty acid esters of ethoxylated fatty alcohols, lanolin and derivatives such as lanolin, lanolin oil, lanolin wax, lanolin alcohols, lanolin fatty acids, isopropyl lanolate, ethoxylated lanolin, ethoxylated lanolin alcohols, ethoxylated cholesterol, propoxylated lanolin alcohols, acetylated lanolin alcohols, lanolin alcohols tinoteate, lanolin alcohols ricinoleate, ethoxylated alcohols-esters, hydrogenated lanolin, ethoxylated hydrogenated lanolin, and ethoxylated sorbitol lanolin; polyhydric alcohol esters such as ethylene glycol mono- and di-fatty acid esters, diethylene glycol mono- and di-fatty acid esters, polyethylene glycol (200-600) mono- and di-fatty acid esters, propylene glycol mono- and di-fatty acid esters, polypropylene glycol 2000 mono-oleate, polypropylene glycol 2000 mono-stearate, ethoxylated propylene glycol monoostearate, glyceryl mono- and di-fatty acid esters, polyglycerol poly-fatty acid esters, ethoxylated glyceryl monostearate, 1,3-butylene glycol monostearate, polyoxyethylene polyot fatty acid ester, sorbitan fatty acid esters, and polyoxyethylene sorbitan fatty acid esters; wax esters such as beeswax, spermaceti, myristyl myristate, and stearyl stearate; beeswax derivatives such as polyoxyethylene sorbitol beeswax; vegetable waxes such as camauba and candelilla waxes; phospholipids such as lecithin and derivatives thereof; and sterols such as cholesterol and cholesterol fatty acid esters. However, emollients based on compounds which contain active amine groups are unsuitable. For example, amides such as fatty acid amides, ethoxylated fatty acid amides, and solid fatty acid alkanolamides should not be used,

Emulsilying Agents including Emulsilying and Stiffening Agents and Emulsion Adjuncts - are used for preparing to oil-in-water resultions forming part of the present invention, e.g., non-lone insulfaties such as Ω_{10} - Ω_{20} and sught abcohols and said fatly alcohols condensed with from 2 to 20 moles of ethylene oxide or propylene oxide, $(C_{\rm p} C_{12})$ ality in encoordinated with from 2 to 20 moles of ethylene oxide, in-non- and dis- $C_{\rm p} C_{20}$ stay acid estars of einylene glycol, polyethylene glycols of MW 200-6000, polypropylene glycols of MW 200-6000, and particularly sorbitol, sorbitan, polyocyethylene sorbital, polyocyethylene sorbital, hydrophilic wax estrey, decisionally alcohol, clyel achorol, lanoin abcohols, cholesterol, mono- and disperdies, glycoly monostearate, polyethylene glycol monostearate, mixed mono- and distearic esters of ethylene glycol and polyocyethylene glycol monostearate, and hydrocypropylene glycol subsets.

However, emulsifying agents which contain active armine groups should not be used. Such agents typically include annion emulsifiers such as fally act seaps, e.g., actium, potassium and triethandamin seaps of $C_{10}C_{20}$ jail vajoritis, alkali metal, ammonium or substituted ammonium $(C_{10}C_{20})$ jailly fall metal, $C_{10}C_{20}$ jailly suffered to substituted ammonium $(C_{10}C_{20})$ jailly fall metal, ammonium or substituted ammonium $(C_{10}C_{20})$ jailly fall metal, and jailly fall when suffered suffered in the substituted with a substituted and polymers of 2-propencie acid fogother with polymers of acrylic acid, both cross-trinked with ally inhers of sucross and/or pointeryfirthiol, whing yarying viscosities and identified by product names carbonner 910, 934, 9347; 940, 941, or 134. 1342 however, cationic emulsifiers having active amine groups should also be avoided, which would include those based on quaternary ammonium, mepholnium and pyridinium compounds. Similarly, amphoteric emulsifiers having active armine groups, such as coorbalanes, launy dimethylarine oxide and coorginitizations, should not be used. Useful emulsifying and stiffening agents includedly abord and sodium stearate, and emulsion adjuncts such as olais acid, stearie acid, and sleany ladochol.

Excipients - include, e.g., laurocapram and polyethylene glycol monomethyl ether.

"Humestants - are compounds which promote retention of moisture, thus often being referred to as moisturizers, and include, ag., scroticel, gybranin, glycereth 7 triscateate, glycereth 7 discanancet, hexanetriol, hexylene glycol, propylene glycol, alkoxylated glucose, D-panthenol and derivatives thereol, and hyairunce acid. However, as with the other above-discussed additives, humestants teving active amine groups should not be used, and those include, 6, g., lactamide monesthenolamine and acetamide monoethanolamine. Propliene oxide ethers of gybranin and hydrolyzed corn starch are suitable for use, but humestants based on vegetable protein are not, and these include hydrolyzed wheat protein/wheat oligosactoriaries, hydrolyzed comp protein hydrolyzed wheat protein hydrolyzed soy protein, hydrolyzed rice protein, and hydrolyzed potato crotein.

Ointment Bases - include, e.g., petrolatium, polyethylene glycol, lanciin, and potoxamer, which is a block copolymer of polyoxyethylene and potyoxypropylene which may also serve as a surfactant or emulsifying agent, and which may be represented by the following structural formula:

/XVI

Penetration Enhancers - include, e.g., dimethyl isosorbide, diethyl-glycol-monoethylether, 1-dodecylazacycloheptan-2-one, and dimethylsulfoxide (DMSO).

Perfumes, Fracrances and Other Aesthetic Components: include, 8 g., peppermint, rose oil, rose water, aloe vera, clove oil, menthyl, carphor, deucalybus oil, ougend, menthyl lactais, witch hazel distillate, allantion, bisabold, dipotensium of the properties of the

<u>Preservatives</u> - are used to protect the pharmaceutical or cosmelological composition from degradative attack by, ambient microorganisms, and include, e.g., benzalikonium chloride, benzethonium chloride, alkyl esters of p-hydroxybenzoic acid, lydacthio derivatives, calybylyridinium chloride, monohioghyceru, þrendy, þendryvelhanol, methylparagen, imidazolidinyl urea, sodium dehydroacetate, propylparaben, quaternary ammonium compounds, especially polymers such as polixetonium chloride, potassium benzoste, sodium formaldehyde sulfoxylate, sodium propionate, and thirmerosal.

Sequestering Agents - comprising cyclodextrins have been used heretofore to improve the stability of dihydroxyacetone and other active agents. The cyclodextrins are a lamily of astural cyclic oligosaccharides capable of forming inclusion complexes with a variety of materials, and are of varying ring sizes, those having 6, 7-and 8-gluccas residues in a ring being commonly referred to as c-cyclodextrins, 8-by-clodextrins, 8-cyclodextrins, especially sizes, the procedextrins includes, 9,9, c-cyclodextrin, 8-cyclodextrin, 8-by-clodextrin and calionized cyclodextrins.

Solvents - a number of which have already been discussed, include, e.g., acetone, alcohol, amyene hydrate, butyl alcohol, com oil, cottonseed oil, ethyl acetate, glyconn, hexylene glycol, isopropyl alcohol, isostearyl alcohol, methyle alcohol, methylene chloride, mineral oil, peanut oil, phosphoric acid, polyethylene glycol, polyoxypropylene 15 stearyl ether, propylene glycol, propylene glycol discotate, sesame oil, and purified water.

Stabilizers - include, e.g., calcium saccharate and thymol.

Stiffening Agents - include, e.g., cetyl esters wax, myristyl alcohol, parafin, synthetic parafin, emulsifying wax, microcrystalline wax, white wax and yellow wax.

Sugars - are sometimes used with self-taining compositions to improve the results obtained, and include, e.g., monosaccharides, disaccharides and polysaccharides such as glucose, xylose, fructose, reose, ribose, pentose, arabinose, allose, tallose, altrose, mannose, galactose, lactose, sucrose, erythrose, glyceraldehyde, or any combination thereof

Sunscirem Agents: which are conventionally employed to block or reduce the amount of ultravoler radiation impinging upon the skin, and also be used in combination with the self saming compositions of the present invention; as an adjunct or as complementary freetin, aller other and are difficial tan to be achieved at the same time. Typical sunscired agents include both organization and regimental tan to be achieved at the same time. Typical sunscired agents include both organization from ultravoler tradiation by mechanism which include absorption, and are generally surmised to precise tradiation from ultravoler tradiation by mechanism which include absorption, in the air may be employed for stemple and the skin and the same time and the sunscired agents well known in the air may be employed for stemple purposes in the compositions of the present invention and include, e.g., 2 estim-havyl-p-methoxycrimanisms of the present invention and include, e.g., 2 estim-havyl-p-methoxycrimanisms and the stemple purposes in the compositions of the present invention and include, e.g., 2 estim-havyl-p-methoxycrimanisms and the stemple purposes in the compositions of the present invention and include, e.g., 2 estim-havyl-p-methoxycrimanisms and the stemple purposes in the compositions of the present invention and include, e.g., 2 estim-havyl-present purposes. In the composition of the present invention and include, e.g., 2 estim-havyl-present invention and include, e.g., 2 estim-havyl-present invention and include asset with 4 expensive purposes. In the composition of the present invention and include asset with 4 expensive purposes.

Surfacions - are employed to provide stability to multicomponent compositions, enhance existing properties of
Surfacions - are employed to provide stability to multicomponent compositions, enhance existing properties of
those compositions, and bestow desirable new characteristics to said compositions. They are used as welling agents,
antifloam agents, for reducing the surface tension of wells, and as emulalifiers, dispensing agents and penetrants, and
include, e.g., lapyrium chloride, laureth et.g., ancitation of postery-poly(cop-tension) exhanced polytopien agents and penetrants, and
include, e.g., lapyrium chloride, laureth et.g., a mixture of postery-polytopien polytopien glycol monocologic ethers averaging about 9 estylene
coxide groups per molecule; monochbanclamine, noncoynel 4, 9 and 10, 1e., polytopiene glycol mono(pony)phenyl)ether, nonexyn of 15, i.e., oc. (pony)phenyl)-bytocy-polytopiene glycol monochbance polytopiene,
approx. 3000, poloxwarer (see oriment bases above), polytoys 8, 40 and 50 seatts (e.g., polytoys-12-ethanedly),
callydroxy-polytopiene). Sorbitan, monochbance above, polytoys 4, 6 and 50 seatts (e.g., polytoys-12-ethanedly),
polytorsy-polytopiene 20, i.e., a contian, included carenous, polytoys-12-ethanedly),
polytorsy-polytopiene 20, i.e., a contian, included carenous, polytoys-12-ethanedly),
polytopiene 20, i.e., sorbitan, monocharacteriscense, polytoys-12-ethanedly),
polytopiene 20, i.e., sorbitan, monoclaurate, sorbitan monocl

monostearia, sorbitan sesquiclearis; sorbitan tricieate; and sorbitan tristearria A further description of additional types of auxiliary ingredients which may optionally be included in the compositions of the present invention, as well as of specific compositions of those types and of the above-described types which may thus be optionally included, may be found in the CTFA international Cosmetic Ingredient Dictionary 4th ed., The Cosmetic, Toilefry, and Fragrance Association, Inc., Washington, D.C., 1991.

Lubricity, which refers to the property of having a smooth or slippen guality, is a desirable property of the compositions of the present invention that can be enhanced by the addition of agents well known in the aff of this purpose. In particular, a silicone oil or fluid is preferably used since it can provide the desired lubricity and emollience without a greasy feet. Examples of such agents include dimethyl polysicoune and methylphenyl polysicoune which are water, soluble, and silicone glycol copolymer which is alcohol soluble. Polysicounes with or are commonly employed include dimethyl polysicoune which is and-blocked with trimethyl units, the CTFA name for which is dimethicone, and poly-dimethylycicolosiume, the CTFA name for which is cyclomethicone. Such preferred silvasines solubit at viscosity of from about 2 to about 50 centistokes at 25° C, and are employed in sufficient amount to aid in stimulating removal of other scale from the skin and to aid in controlling conditioning of the skin, which will usually be from about 1.0% to about 1.0.0% by weight, based on the total weight of the composition, and preferably will be from about 1.0% to about 3.0% by weight.

The cosmitologically acceptable carrier employed in the compositions of the present invention may be an emulsion which is a allicone-in-water emulsion. Water-in-cil emulsions can also be utilized to prepare jotions and creams, and oil in that event will be the continuous phase, and the ratio of the amounts of oil to water will be in the range of from about 2: 10 about 1:100, a silicone-in-water amulsion is typically comprised of a mixture of hydrocarbons and silicones in water, a g., a matture of hydrocarbons and silicones in water, a g., a matture of hydrocarbons, volatile allicones, and allystated derivatives of polymeric silicones such as hydrogenated polystobutene, cyclomethicone, and catyl crine-thorn. The term "votalite" as applied to silicono lict, refers to those materials which have a measurable vapor pressure at ambient temporatures. Votalite silicone oils are preferably chosen from cyclic or linear polymerity isliconaes containing from about 3: 0 about 9, and preferably from about 4: 0 about 5, alliconae in temporatures. The nonvotalite discinous oils evitor and a useful unit preparature copylimers. Include polymers allicone copylimers include a polywarplaylene other copylimer having a viscosity of about 1200 to 1500 centistokes at 25°C, available as SF-1066 organosilicone sur-catant from General Electric. Celly dimethicone copolymer indemicricone are specially useful because they also function as emulalities and emolliants. In such compositions, as well as in other compositions of the present invention. It has been found that to sclutum chorido is a usualist stabilizing agent.

Indeed, the overall viscosity of the compositions of the present invention may exhibit considerable variation, some of which is by design and is a product of the type of formulation which is being prepared. For example, a cream formulation will inherently have a higher viscosity than an aerosol spray formulation. Generally, however, with the two key properties of stability and dispersibility being of paramount interest, the viscosity of the compositions of the present invention will desirably between about 2000 and about 2000 Cgn bern the viscosity is below about 2000 cps., the compositions will exhibit a reduced stability, whereas, when the viscosity is above about 20,000 cps., it will be difficult to disperse the compositions of the present invention by ordinary means.

The compositions of the present invention may be prepared using very straightforward methodology which is well understood by the arisand or oftendy skill. Where the compositions of the present invention are simple aqueous and/or other solvent solutions, the various components of the overall composition are brought together in any practical roder, which will be dictated largely by considerations of convenience. Those components having reduced water sofulbility, but sufficient solubility in the same oc-solvent with water, may all be dissolved in as add co-solvent, after which the co-solvent solution will be added to the water portion of the carrier whereupon the solutios therein will become dissolved in the water. To add in this dispersion/solution process, a sufficient and water in the prophysic dispersion of the carrier whereupon the solution will be come dissolved in the water. To add in this dispersion/solution process, a sufficient and water.

Where the compositions of the present invention are to be in the form of emulsions, as where the resultant formulation is a billion of a ordinary the components of the composition will be brought foundation as which or a ordinary the components of the composition will be brought foundation as the following general procedures. The continuous water phases is first heated to a temperature in the range of from about 50° to about 95° C., preferably from about 75° to about 85° C., preferably from about 80° C., the choice of which temperature will depend, of course, upon the physical and chemical properties of the components within make up the oil-in-vater armission. Once the continuous water phases has reached its selected temperature, the components of the composition which are not be adoled at this stage, referred to a Phase 1, are admixed with the water and dispersed therein under high-speed agitation. Next, the temperature of the water is restored to appoximately its original level, after which the Phase II components of the composition and mixing continues for from about 50 about 60 minutes, preferably about 10 to about 30 minutes, depending on the components of the composition without parties of thom about 20° to about 50°C., preferably to from about 35° to about 45°C., after which the composition mixture is a retirely colorated from about 20° to about 50°C., preferably to from about 35° to about 45°C., after which the composition of the or arborning of the asset in the acceptance of passively or passively or passively or passively

cooled to from about 20° to about 40°C., preferably to about 30°C., after which the fragrance phase is added to the composition mixture, to which water is then added in sufficient quantity to reach its original predetermined concentration in the overall Composition.

In accordance with the present invention it has been discovered that substantially improved chemical, including instability, are be conferred upon aqueous formulations containing self stanning skin coloring agents subject to such instability, especially where the self-tanning agent is an en-lydroxy aldehyde or ketone, more especially 1,3-dihydroxy-acetone. The basic mechanisms whereby such self-tanning agents are stabilized in accordance with the present invention are contemplated to be applicable across a horsed ranged of such estimating agents. Accordingly, the discussion which follows, relating to such mechanisms of action which are hypothesized to take place with regard to the stabilization of aqueous formulations containing 1,3-dihydroxyscatone, will be understood to be merely demonstrative of the present invention, and therefore to be applicable to many other self-tanning agents within the scope of the present invention.

This compositions of the present Invention utilize water-soluble alcohole ("clot"), alkoxylated ols and dhydric alcohols ("clote") together to provide chomical and pt stability for up of a house at 80° c, which is comparatively much superior to the results obtained with aqueous formulations which do not contain these components. It is theretized that the keto functionality of 1,3-dihydroxysection is capable or therefore the termation, analogous to acatal formation, an the presence of 1,2-disk and 1-pt. ("and that the use of 1,2-disk or polypic containing vicinal hydroxyl functions, at a 1:1 molar ratio, is required for this chemical pathway to operate efficiently, it is conjectured that the presence of H₂O" in this system shifts the equilibrium of the reaction in this pathway from a keto-containing molety to a five-membered cyclic ketal structure which is more stable than the monomeric 1,3-dihydroxysection-(orm and thus more resistant to degradation. This theorized pathway may be prepresented by the following reaction:

It is thought that within a pH tange of form 3.5 to 4.5 and at antibient importatures, 1,3-dihydroxyscitone is in aquious formulation within the scope not he present invention is applied to the skin, the pH of the system is elevated to 5.5 which substantially reverses ead resection, 1.6, nation expected to the skin, the pH of the system is elevated to 5.5 which substantially reverses ead resection, 1.6, nation expected to the skin, the pH of the system is elevated to 5.5 which substantially reverses ead resection, 1.6, nation expected formation. There is produced on the skin as a formation there is no standard resection with the skin to take place, which results in the desired artificial tarning, it may be seen, accordingly, that the present invention provides a means for stabilizing 1,3-dihydroxyscetone for long periods of time by conventing into a cyclic ketal while it is being stored as an acquious termination ready for use. The cyclic ketal is stable but incepable of producing the self-tanning reaction on the skin. Surprisingly, this same stabilized aqueous formulation of the present invention, when applied to the skin, permits reversion of the cyclic ketal back to the unstable 1,3-dihydroxyscetone, which applied produces the self-tanning reaction, making its ansatiality of no consequence.

It is further theorized that the keto functionality of 1,3-dihydroxyscatone undergoes cyclic ketal formation in the presence of polyois ("clos") and H₂O" and that the use of such a water soluble alcohor in a 21 mater ratio is required for this pathway to operate efficiently. Although 1,3-dihydroxyscatone reduces the pH of an aqueous solution to which it is added, the presence of H₂O" produces stability in this system by shifting the equilibrium of the reaction in this pathway from a keto-containing mixely to a more stable five-membered cyclic ketal structure, as already above-described. This pathway may be represented by the following scheme:

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With some polyols, it is possible to form more than one cyclic ketal with the polyol, as is the case with mannion as illustrated in the following reaction scheme:

An alcohol which has demonstrated good synergy with 1,3-dihydroxyacetone is benzyl alcohol. Although benzyl alcohol is commonly used as a preservative in cosmetologic products, it is theorized that this aromatic alcohol reacts with 1,3-dihydroxyac etone to form a more stable cyclic ketal in much the same manner as above described with respect to short chain primary alcohols, but at a much slower rate of reaction. The phenyl functionality of benzyl alcohol confers upon it a degree of hydrophobicity which makes it impracticable to use 2:1 molar ratios of the benzyl alcohol, its solubility in water being exceeded at these levels.

It is further theorized that hydrogen bonding plays an important role in the stabilization of 1,3-dihydroxyacetone in accordance with the present invention. In aqueous solution, 1,3-dihydroxyacetone is converted from a dimeric to a monomeric form, only the latter being active as a self-tanning agent. Hydrogen bonding plays a role in maintaining 1,3-dihydroxyacetone in the monomeric form, and if sufficient portion of the aqueous solvent are bound by other excipients, or are lost by evaporation, the amount of possible hydrogen bonding is reduced.

It is thought that the -ols, -diols, and alkoxylated ethers present in the compositions of the present invention aid in providing sufficient hydrogen bonding at the hydroxyl functionalities of 1,3-dihydroxyacetone well solvated and thus in the desired monomeric form. Alkoxylated ethers, especially diethylene glycol monoethyl ether, i.e., ethoxydiglycol, exhibit acidity in aqueous solution which will be favorable to the conditions necessary for optimal 1,3-dihydroxyacetone stability. Ethoxydiglycol, present in the amounts preferred for the compositions of the present invention, will result in a pH of said composition of about 4.7, pnor to addition of the 1,3-dihydroxyacetone. This factor, in addition to the three site available for hydrogen bonding, makes ethoxydiolycol a preferred component for the compositions of the present invention.

DESCRIPTION OF PREFERRED EMBODIMENTS

There follows a description of certain preferred embodiments of the compositions of the present invention, as well as a description of the methodology employed to evaluate the improved stability and other enhanced properties of said compositions, including the results of those evaluations. However, the following description is presented solely for the purpose of further illustrating the invention and is in no way intended to be, nor should it be construed to be a limitation of any sort on the scope or any of the essential or auxiliary features of the present invention, which are to be determined and defined solely by the attendant claims which form a part hereof.

EXAMPLE 1

Self-Tanning Cosmetic Formulation in Spray Form for Producing a Dark Tan

The individual components of the spray formulation, identified in Table 1 below, were brought together in four phases, i.e., four different groups of components were added separately to the vessel in which the formulation was being prepared largely as a matter of convenience. As a practical matter, all of the components of the formulation were brought into admixture with each other in a single step, and the order of addition was not critical, but was, rather, also a matter of convenience.

TABLE 1

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COMP.
1.1	Deionized Water		70.75
1	Edetate Disodium	USP1	0.10
1 :	Ethoxydiglycol		20.00
1 Tag	Benzyl Alcohol (and) Methyl-paraben (and) Propylparaben	× -	0.35
8 1 39	Sorbitol Solution	USP	1.00
* -1	Dihydroxyacetone		4.00
. 11	Nonoxynol-9	•••	0.50
	Fragrance		0.10
111	Panthenol		0.20
. 111	Retinyl Palmitate, Water Miscible		0.20
10	Propylene Glycol (and) Walnut Extract		1.00
III	Butylene Glycol (and) Water (and) Alce Extract (and) Matricaria Extract (and) Conellower Extract (and) Ouillaja Saponaria Extract (and) Witch Hazel	-	1.50
11	Extract	100	
	Propylene Glycol (and) Water (and) Kiwi Fruit Extract	\$ 6.76 h	0.20
IV .	Sorbic Acid	NF2	0.05
IV	Sodium Benzoate	NF	1 - 14 - 14 - 10.05

United States Pharmacopeia, Rockville, MD 2085

EXAMPLE 2

Self-Tanning Cosmetic Formulation in Spray Form for Producing a Deep Dark Tan

The individual components of the spray formulation, identified in Table 2 below, were brought together in essentially the same manner as above-described in Example 1. Four phases of components were added separately to the vessel, largely as a matter of convenience. The darker has produced by the formulation of this example, compared to that produced by the formulation of Example 1 above, is the result of 50% more of the active self-tenning agent, dhydroxy-actions, present in this example.

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COMP.
1	Deionized Water	1	68.75
	Edetate Disodium	USP	0.10
	Ethoxydiglycol	2	20.00
j i s	Benzyl Alcohol (and) Methylparaben (and) Propylparaben		0.35
1 0	Sorbitol Solution	USP -	1.00
1 11	Dibudroxyacetone		6.00

² National Formulary

ED O OOA OAE AA

TABLE 2 (continued)

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COMP.
. 11	Nonoxynol-9	7	0.50
. # -	Fragrance		²⁻ - 0.10
10	Panthenol -		0.20
10 ° 10	Retinyl Palmitate, Water Miscible		0.20
101	Propylene Glycol (and) Walnut Extract		1.00
=	Butylene Glycol (and) Water (and) Aloe Extract (and) Matricana Extract (and) Coneflower Extract (and) Quillaja Saponana Extract (and) Witch Hazel Extract	i ∓3	1.50
=	Propylene Glycol (and) Water (and) Kiwi Fruit Extract	N.T.; 4.	0.20
IV	Sorbic Acid	NF	0.05
IV	Sodium Benzoate	NF	0.05

EXAMPLE

Self-Tanning Cosmetic Formulation in Oil-in-Water Emulsion Cream Form for Producing a Deep Dark Tan

The continuous water phase was first heated to a temperature of 75° io 80°C, after which the components of Phase I set out in Table 3 balow, were admixed with the water and dispersed therein under high-speed agitation. Next, the temperature of the water was restored to 75°C, tater which the Phase II components as do ut in Table 3 were added to the composition mixture under moderate agitation, and mixing continued for 10 to 30 minutes. Thereafter, the composition mixture was actively cooled to 35° to 45°C, after which the components of the remaining Phases III, IV and V set out in Table 3 were combined and added to the composition mixture. The composition mixture was then actively cooled to 30°C, after which the fragrance phase was added to the composition mixture, to which water was then added in sufficient quantity to reach the concentration set out in Table 3.

	25	100	100	16 22	 T 6	TARI

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COMP.
1	Purified Water	USP	67.800
143.9	Xanthan Gum	NF	0.350
1.4	Ethoxydiglycol	·	5.000
i, de, 4	C ₁₀ -C ₃₀ acrylate cross-polymers ³		0.100
$f \in \Gamma_{i}^{1}$	Sorbitol 70% By Weight Solution	NC4	5.000
	Diethanolamine cetyl phosphate		0.100
1117	Cetyl Alcohol	NF	1.500
- JI '	Stearyl Alcohol		1.500
.113	Steareth-20		1.000
- 11	Octyl Palmitate		1.500
11	Dimethicone Copolyol ⁵		1.000

Non-crystallized = USP grade.

⁵ Available from Goldschmidt under the trademark Abit Way 9001

⁶ Polyethylene glycol, approx. mol. wt. of 100.

TABLE 3 (continued)

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COME
11 - 1	Glyceryl Stearate (and) PEG-1006Stearate7		0.500
11.	α-Tocopherol Acetate (Vitamin E)		0.100
- 0	Dimethicone 3508		0.050
0 H 5	Actiplex 335 Lipo M		0.100
H 550	Glucan P-20 Distearate9		1.500
· II , 5	Dihydroxyacetone	/ ··· + = 3	5.000
111	Plant Extracts ¹⁰	E	0.250
IV	Sodium meta-Bisulfite	, ,-	0.025
IV -	Purified Water	USP	6.150
IV	50% Aqueous DL-Panthenol		0.400
. V	Stearyl Glycerrhetinate		0.025
Va	Fragrance		0.100

Polyethylene glycol, approx. mol. wt. of 100

EXAMPLE 4

Evaluation of Stability of Dihydroxyacetone in a Composition of the Present Invention When Subjected to Harsh Storage Conditions

In order to evaluate the extent to which a typical composition of the present invention was able to protect the dihydroxyacotions (DHA) contained in said composition from degradation under harin storage conditions, the composition set out in Table 5 below was stored at 25°C. for three (3) months and at 5°C. for three (3) months

TABLE

COMPOSITION STORAGE CONDITIONS	BEGINNING pH	FINAL pH	BEGINNING DHA CONC. (wt. %)	FINAL DHA CONC. (wt. %)
3 mos. at 25°C.	4.7	4.3		and the state
2 mos. at 50°C.	4.7	4.0	6.0	5.2

TADLES

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COMP.
el de	Purified Water	USP	68.75
3 1 1	Edetate Disodium	USP	0.10
11	Tivalin SF		20.00
7	Plant Extracts ¹¹		0.35
	Sorbitol Solution	USP	1.00

TI Extracts of explic elects available from Nion under the trademark Nipaguard MP

⁷ Available from under the trademark Artacel 165.

^{8 350} centistokes (cst).

Polypropylene-20/glucan distearate ether.
 To Extracts of exotic plants evallable from Nipa under the trademark Nipagus

TABLE 5 (continued

PHASE NO.	INGREDIENT DESCRIPTION	GRADE	% BY WT. PER TOTAL WT. OF COMP
, I .	Dihydroxyacetone		6.00
II	Nonoxynol-9		0.50
JI 1 2	Fragrance		0.10
lii .	dL-Panthenol		0.20
- III 💎	Retinyl Palmitate, Water Miscible		0.20
- 111	Propylene Glycol (and) Walnut Extract		1.00
=	Butylene Glycol (and) Water (and) Aloe Extract (and) Matricaria Extract (and) Conellower Extract (and) Quillaja Saponaria Extract (and) Witch Hazel Extract	70 V	1,50
1 (III - 3) 1 (1)	Propylene Glycol (and) Water (and) Kiwi Fruit Extract		0.20
IV ·	Sorbic Acid	NF	0.05
IV	Sodium Benzoate	NF ·	0.05

Claims

1. - A composition comprising:

- (A) from about 0.5% to about 20.0% by weight, based on total weight of said composition, of a self-tanning skin coloring agent subject to chemical instability;
- (B) from about 2,0% to about 40,0% by weight, based on total weight of said composition of a polyethoxyglycol;
- and
 (C) from about 0.1% to about 15.0% by weight, based on total weight of said composition, of a polyol comprising
 a polyhydric compound having at least three hydroxyl groups and at least three carbon atoms.
- A composition according to Claim 1 wherein said self-lanning skin coloring agent subject to chemical instability is an α-hydroxy aldehyde or ketone.
 - A composition according to Claim 2 wherein said α-hydroxy aldehyde or ketone is dihydroxyacetone, and is present in an amount of from about 4.0% to about 6.0% by weight, based on the total weight of said composition.
 - A composition according to Claim 1 wherein said polyethoxyglycol is ethoxydiglycol, and is present in an amount
 of from about 14.0% to about 25.0% by weight, based on the total weight of said composition.
- 5. A composition according to Claim 1 wherein said polyol comprises one or more members independently selected from the group consisting of 1,2,6-hexanetriol, isopropylidene glycerol, polyocyethylene sorbiols, glycerin (glycerol), eighylorin, eytyhribol, mannitol, yritiol, D and L-sorbiol, glycese, furcles, glaaclose, mannose, sucrose, lactose, triebalse, mallose and inositol, and said polyol is preferably D-sorbiol, D-mannitol, or inositol.
- A composition according to Claim 5 wherein said polyol is D-sorbitol, and is present in an amount of from about 0.5% to about 1.5% by weight, based on the total weight of said composition.

7. A composition comprising:

- (A) from about 0.5% to about 20.0% by weight, based on total weight of said composition, of a self-tanning skin coloring agent subject to chemical instability;
- (B) from about 2.0% to about 40.0% by weight, based on total weight of said composition of a polyethoxyglycol, preferably ethoxydiglycol;
- (C) from about 0.1% to about 15.0% by weight, based on total weight of said composition, of a polyol comprising

a polyhydric compound having at least three hydroxyl groups and at least three carbon atoms, and (D) from about 0.1% to about 8.0% by weight, based on total weight of said composition, of a water soluble dihydroxyl compound having at least two, and up to eight carbon atoms.

- A composition according to Claim 7 wherein said dihydroxyl compound has from two to five carbon atoms and is
 present in an amount of from about 0.4% to about 2.0% by weight, based on total weight of said composition.
 - 9. A composition seconding to Claim 9 wherein said self-tening skin coloring agent subject to chemical instability is an a-hydroxy aidenlyde or ketone comprising dihydroxyacatone present in an amount of from about 4.0% to about 6.0% by weight, based on the lotal weight of said composition; said polyethoxyglycol is ethoxydiglycol present in an amount of from about 14.0% to about 25.0% by weight, based on the Iclaid weight of said composition; and said polyor compress on our ornor members independently selected from the group consisting of 1,26-hearantirol, isopropylidene glycerol, polyoryethylene sorbitols, glycerin (glycerol), diglycerin, erythrifo, manniot, xyllal, D and L-sorbitol, glycers (putoses futoses, gliatoses, surrose, surrose, surrose, lostose, threakboer, midstes and nostal or.
 - 10. A composition according to Claim 8 wherein said dihydroxyl compound comprises one or more members selected mit be group consisting of ethylene glycot, diethylene glycot, triethylene glycot, triethylene glycot, ether soluble polywrithylene glycot, ethylene glycot, ethylen
 - A composition according to Claim 8 wherein said polyol is D-sorbitol, D-mannitol, or inositol present in an amount
 of from about 0.5% to about 1.5% by weight, based on the total weight of said composition.
- 25 12. A composition comprising:
 - (A) from about 0.5% to about 20.0% by weight, based on total weight of said composition, of a self-tanning skin coloring agent subject to chemical instability.
 - (8) from about 2.0% to about 40.0% by weight, based on total weight of said composition of a polyethoxyglycol, preferably ethoxydiglycol;
 - (C) from about 0.1% to about 15.0% by weight, based on total weight of said composition, of a polyol comprising a polyhydric compound having at least three hydroxyl groups and at least three carbon atoms;
 - (D) from about 0.1% to about 8.0% by weight, based on total weight of said composition of a water soluble
 - dihydroxyl compound having at least two, and up to eight carbon atoms; and

 (E) an additying agent in amount sufficient to maintain the pH of said total composition at from about 3.5 to
 - about 4.5.
- 13. A composition according to Claim 12 wherein said pH of said total composition is about 4.0, and said acidfying agent is one or more members selected from the group consisting essentially of acetic, adipic, anisic, benzoic, boric, carbonic, cinnamic, citric, diphosphoic, formic, tumaric gallic, glutaric, glycocis, factic, maleic, maleic, maleic, maleic, maleic, maleic, priballic, propositic, pyruvic, salicytic, succinic, tartaric and vanillic acid.
 - 14. A composition according to Claim 13 wherein said additying agent is selected from the group consisting essentially of sorbic acid, acetic acid, maleic acid, cliric acid and furnaric acid, present in an amount of from about 0.01% to about 0.05% by weight, based on the total weight of said composition.
 - 15. A composition according to Claim 2 comprising/rom about 0.5% to about 20.0% by weight, based on total weight of said composition, of dihydroxyacetone of the formula:

(1)

16. A composition according to Claim 4 comprising from about 2.0% to about 40.0% by weight, based on total weight of said composition of ethoxydighycol of the formula:

(III-a)

- 17. A cosmetologic product for application to the hair, nails or skin of a subject for the purpose of tanning, coloring and/or darkening the same, comprising a composition according to Claim1 wherein a remaining weight percent portion of said composition comprises a cosmetologically acceptable carrier.
- 18. A cosmetologic product according to Ctaim 17 wherein said cosmetologically acceptable carrier comprises one or more members independently selected from the group consisting of actifying and alliatizing agents, aerosol propellants, antimicrobial apents including antibacteral, antitungal and antiprotozoal agents, antimicrobial preservatives, antioxidants, buffering agents, clothating agents, clothing additives including dyes and pigments: demaid-logically active agents, depersing agents, smollients, emulsiying agents including emulsiting and stiffening agents and emulsion adjuncts; excipents; humeclants; onthent bases, penetration enhancers, perfumes and fragrances; preservatives; exceptating agents, solvents, stabilizes; sulgens; supers, supe
- 19. A method of tanning, coloring or darkening the hair, nails and/or skin of a subject comprising applying thereto an amount of cosmetologic product according to Claim 17 sufficient to tan, color, or darken said hair, nails and/or skin of said subject to which it is applied to the extent desired by said applicant.
- 25 20. A method for preparing a cosmetologic product according to Claim 17 wherein said product is an aqueous solution, comprising the following steps, carried out sequentially or simultaneously:
 - (1) combining the following components of said product: water, a self-tanning skin coloring agent, a polyethory sylpcol and a polyol, opticnally together with, chiefsing agents, sequestering agents, and antimicrobial preservatives, and/or antioxidants which are desired, and optionally with a solvent therefor, and thereafter optionally. (2) acting a perturne or tragrames which is desired optionally with a solvent therefor.
 - (3) adding vitamins, nutrients, penetration enhancing agents, coloring additives, sunscreen agents, and/or dermatologically active agents which are desired, optionally with a solvent therefor, and
 - (4) adding antimicrobial preservatives which are desired, optionally with a solvent therefor.
 - 21. A composition according to Claim 1 comprising:
 - (A) from about 0.5% to about 20.0% by weight, based on total weight of said composition, of an α-hydroxyaldehyde or ketone of the formula



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- \mathbb{R}^1 is H, CH₂OH, CHOHCH₂OH, CH(OH)CH(=O), CH(OCH₃)CH(=O), CH(NH₂)CH(=O), or CH(NH-Pheny)(CH(=O)); and \mathbb{R}^2 is H or CH₂OH. and
- (B) from about 2.0% to about 40.0% by weight, based on total weight of said composition of an polyethoxyglycol of the formula.

(III)

wherein

- n is an integer of from 2 to 6; and
- R³ is H, (C₁-C₆)alkyl, or phenyl.

22. A composition comprising:

- (A) from about 4.0% to about 6.0% by weight, based on the total weight of said composition, of dihydroxyac-
- (B) from about 14.0% to about 25.0% by weight, based on the total weight of said composition of ethoxydigtycol;
- (C) from about 0.5% to about 1.5% by weight, based on the total weight of said composition, of D-sorbitol;
 (D) from about 0.4% to about 2.0% by weight, based on the total weight of said composition, of propylene
- glycol and butylene glycol; and
 (E) from about 0.04% to about 0.20% by weight, based on the total weight of said composition, of sorbic acid.
- A cosmetologic product for application to the hair, nalls or skin of a subject for the purpose of tanning, coloring;
 and/or derkening the same, comprising a composition according to Claim 22 wherein a remaining weight percent portion of said composition comprises a cosmetologically acceptable carrier.
 - 24. A cosmetologic product according to Ctaim 29 wherein said cosmetologically acceptable carrier comprises one or more members independently selected from the group consisting of actifying and alkalizing agents; acrosol propellates; antimicrobial agents including antibactorial, antitungel and antiprotocoal agents; antimicrobial preservatives; antioxidants; buffering agents; chelating agents; cobing additives including dyse and pigments; dematic-logically active agents; dispersing agents; monilatint; emulatilying agent suitoding emulsitying and stiflening agents and emulsion; adjuncts; excipients; humectants; ontiment bases; penetration enhancers; perfumes and fregances; preservatives; exceptating agents; sovents; stablicers; stiflening agents, supracreal agents; sufficient suspending agents, thickening agents; vehicles; viscosity-ocreasing agents; watting agents.



European Patent

EUROPEAN SEARCH REPOR

EP 98 30 4273

	DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document with indication, where appropriate,	Relevant	CLASSIFICATION OF THE
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55.	* claims 1-3,6,7 *-		100
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1	* claims 1,8-10 *	1	110
	* page 10, line 23-25 * * page 11, line 5 - page 12, line 16 *		
911	* example 1 *		
.х	& US 5 514 367 A	A 45.	
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1	* column 4, line 12 - column 5, line 47 *	17.10	
	* column 6, line 34-37 * * examples 1.2 *	11	A61K
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To

WO 2005/077327 PCT/FP2005/050134

From the INTERNATIONAL BUREAU

SECOND AND SUPPLEMENTARY NOTICE INFORMING THE APPLICANT OF THE COMMUNICATION OF THE INTERNATIONAL APPLICATION (TO DESIGNATED OFFICES WHICH APPLY THE 30 MONTH TIME LIMIT UNDER ARTICLE 22(1))

(PCT Rule 47.1(c))

Date of mailing (day/manth/year) 15 June 2006 (15.06.2006)

Applicant's or agent's file reference Rswk-02034

BEIERSDORF AG Unnastrasse 48 20245 Hamburg ALLEMAGNE

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2 0. Juni 2006

FINGANG Kst 6713

IMPORTANT NOTICE

International application No. PCT/EP2005/050134

International filing date (day/month/year) 13 January 2005 (13.01.2005) Priority date (day/manth/year) 11 February 2004 (11.02.2004)

Applicant

BEIERSDORF AG et al

- ATTENTION: For any designated Office(s), for which the time limit under Article 22(1), as in force from 1 April 2002 (30 months from the priority date), does not apply, please see Form PCT/IB/308(First Notice) issued previously.
- 2. Notice is hereby given that the following designated Office(s), for which the time limit under Article 22(1), as in force from 1 April 2002. does apply, hashave requested that the communication of the international application, as provided for in Article 20, be effected under Rule 93bis. 1. The International Bureau has effected that communication on the date indicated below: 25 August 2005 (25.08.2005)
 - AU. AZ. BY. CN. CO. DZ, EP, HU, KG, KP, KR, MD, MK, MZ, NA, PG, RU, SY, TM, US

In accordance with Rule 47.1(c-bis)(i), those Offices will accept the present notice as conclusive evidence that the communication of the international application has duly taken place on the date of mailing indicated above and no copy of the international application is required to be furnished by the applicant to the designated office(s).

- 3. The following designated Offices, for which the time limit under Article 22(1), as in force from 1 April 2002, does apply, have not requested, as at the time of mailing of the present notice, that the communication of the international application be effected under Rule 93bis.1
 - AE, AG, AL, AM, AP, AT, BA, BB, BG, BR, BW, BZ, CA, CR, CU, CZ, DE, DK, DM, EA, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, ID, IL, IN, IS, JP, KE, KZ, LC, LK, LR, LS, LT, LV, MA, MG, MN, MW, MX, NI, NO, NZ, OA, OM, PH, PL, PT, RO, SC, SD, SG, SK, SL, SM, TJ, TN, TR, TT, UA, UZ, VC, VN, YU, ZA, ZW

In accordance with Rule 47.1(c-bis)(ii), those Offices accept the present notice as conclusive evidence that the Contracting State for which that Office acts as a designated Office does not require the furnishing, under Article 22, by the applicant of a copy of the international application.

- 4. TIME LIMITS for entry into the national phase
 - For the designated or elected Office(s) listed above, the applicable time limit for entering the national phase will, subject to what is said in the following paragraph, be 30 MONTHS from the priority date.

In practice, time limits other than the 30-month time limit will continue to apply, for various periods of time, in respect of certain of the designated or elected Office(s) listed above. For regular updates on the applicable time limits (30 or 31 months, or other time limit), Office by Office, refer to the PCT Guzene, the PCT Newsletter and the PCT Applicant's Guide, Volume II, National Chapters, all available from WIPO's Internet site, at http://www.wipo.int/pct/en/index.html.

It is the applicant's sole responsibility to monitor all these time limits.

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20. Switzerland

Agnes Wittmann-Regis

Authorized officer

Facsimile No +41 22 740 14 35

Facsimile No.+41 22 338 89 70

From the INTERNATIONAL BUREAU

PCT

NOTIFICATION CONCERNING SUBMISSION OR TRANSMITTAL OF PRIORITY DOCUMENT

BEIERSDORF AG Unnastrasse 48 20245 Hamburg ALLEMAGNE Eingang 6713 -5. Juli 2005

(PCT Administrative Instructions, Section 411)

Date of mailing (day/month/year) 24 June 2005 (24.06.2005)

Applicant's or agent's file reference

Rswk-02034
International application No.
PCT/EP2005/050134

International publication date (day/month/year)

IMPORTANT NOTIFICATION

International filing date (day/month/year)
13 January 2005 (13.01.2005)

Priority date (day/month/year) 11 February 2004 (11.02.2004)

Applicant

BEIERSDORF AG et al

- 1. By means of this Form, which replaces any previously issued notification concerning submission or transmittal of priority documents, the applicant is hereby notified of the date of receipt by the International Bureau of the priority document(s) relating to all earlier application(s) whose priority is claimed. Unless otherwise indicated by the letters "NR; in the right-hand column or by an asterisk appearing next to a date of receipt, the priority-document concerned was submitted or unamnitted to the International Bureau in compliance with Rule 17.1(a) or (b).
- 2. If applicable) The letters "NR" appearing in the right-hand column denote a priority document which, an the date of mailing at this Earm, had not yet been received by the International Bureau under Rule 17.1(a) or (b). Where, under Rule 17.1(a), the priority document must be submitted by the applicant to the receiving Office or the International Bureau, but the applicant fails to submit the priority document within the applicant to eincreation Rule 17.1(a) which provides that no designated Office may directed to Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no designated Office may directed the Rule 17.1(a) which provides that no design the Rule 17.1(a) which provides the Rule 17.1(a) wh
- 3. (If applicable)An asteriak (*) appearing next to a date of receipt, in the right-hand column, denotes a priority document submitted or the International Bureau but not in compliance with Rule 17.1(a) or (b) (the priority document was received after the time limit prescribed in Rule 17.1(a) or the request to prepare and transmit the priority document was received after the time limit prescribed in Rule 17.1(a) or the request to prepare and transmit the priority document was not furnished in compliance with Rule 17.1(a) to (b), the International Bureau will invertheless transmit a copy of the document to the designated Office, for their consideration. In case such a copy is not accepted by the designated Office as the priority document. Rule 17.1(c) provides that no designated Office and ydisegard the priority claim concluded before giving the applicant an opportunity, upon entry into the national phase, to furnish the priority document within a time limit which is reasonable under the circumstances.

 Priority date
 Priority application No.
 Country or regional Office or PCT receiving Office of Priority document
 Date of receipt of priority document

 11 February 2004 (11.02 2004)
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 15 June 2005 (15.08 2005)

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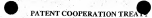
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The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20: Switzerland

Maria KIRCHNER (Fax 338 8970)

Facsimile No. (41-22) 338.89.70 Telephone No. +41 22 338 8056



From the INTERNATIONAL BUREAU

BEJERSDORF AG

Unnastrasse 48

20245 Hamburg

ALLEMAGNE

FIRST NOTICE INFORMING THE APPLICANT OF THE COMMUNICATION OF THE INTERNATIONAL APPLICATION (TO DESIGNATED OFFICES WHICH I DO NOT APPLY THE 30 MONTH TIME LIMIT UNDER ARTICLE 22(1))

(PCT Rule 47.1(c))

Date of mailing (day/month/year) 15 September 2005 (15.09.2005)

Applicant's or agent's file reference

Rswk-02034

International application No. PCT/FP2005/050134

International filing date (day/month/year) 13 January 2005 (13.01.2005) IMPORTANT NOTICE

Priority date (day/month/year) 11 February 2004 (11.02.2004)

Applicant

BEIERSDORE AG et al

- ATTENTION: For any designated Office(s), for which the time limit under Article 22(1), as in force from 1 April 2002 (30 months from the priority date), does apply, please see Form PCT/IB/308(Second and Supplementary Notice) (to be issued promptly after the expiration of 28 months from the priority date).
- Notice is hereby given that the following designated Office(s), for which the time limit under Article 22(1), as in force from 1 April 2002. does not apply, has/have requested that the communication of the international application, as provided for in Article 20, be effected under Rule 93bis. 1. The International Bureau has effected that communication on the date indicated below: 25 August 2005 (25.08.2005)

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In accordance with Rule 47.1(c-bis)(i), those Offices will accept the present notice as conclusive evidence that the communication of the international application has duly taken place on the date of mailing indicated above and no copy of the international application is required to be furnished by the applicant to the designated Office(s).

The following designated Offices, for which the time limit under Article 22(1), as in force from 1 April 2002, does not apply, have not requested, as at the time of mailing of the present notice, that the communication of the international application be effected under Rule 93b(s.):

In accordance with Rule 47.1(c-bis)(ii), those Offices accept the present notice as conclusive evidence that the Contracting State for which that Office acts as a designated Office does not require the furnishing, under Article 22, by the applicant of a copy of the international application

4. TIME LIMITS for entry into the national phase

For the designated Office(s) listed above, and unless a demand for international preliminary examination has been filed before the expiration of 19 months from the priority date (see Article 39(1)), the applicable time limit for entering the national phase will, subject to what is said in the following paragraph, be 20 MONTHS from the priority date:

In practice, time limits other than the 20-month time limit will continue to apply, for various periods of time, in respect of certain of the designated Offices listed above. For regular updates on the applicable time limits (20 or 21 months, or other time limit), Office by Office, refer to the PCT Gazette, the PCT Newsletter and the PCT Applicant's Guide, Volume II. National Chapters, all available from WIPO's Internet site, at http://www.wipo.int/pct/en/index.html.

It is the applicant's sole responsibility to monitor all these time limits.

The International Bureau of WIPO 34, chemin des Colombettes

1211 Geneva 20, Switzerland

Authorized officer

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